

- 32. Vasodilator delivery systems specially adapted to deliver about 5 to 25% of conventional dosage of vasodilators and marked with the appropriate DRG and/or ICD disease codes and/or instructions for titrating or tapering their use, to facilitate their proper application for treatment of diseases involving vasospasm.
- 33. A delivery system according to Claim 32 adapted for transdermal delivery.
- 34. A delivery system according to Claim 32 adapted for delivery of about 0.02 to 20 milligrams per day (Nitroglycerin equivalent) of vasodilator.

35. A delivery system according to Claim 32 adapted for delivery of about

- 0.02 to 20 milligrams per day (Nitroglycerin equivalent) of a vasodilator selected from the group comprising Nitroglycerin in pill, patch, ointment, cream, inhaler, spray and other forms, Nitroglycerin equivalents and substitutes, comprising p.o. clonidine, Dynacire (isradipine), hydrazine, nifedipine, and/or other medicines selected from the empirical group of medications which have the common characteristic of causing smooth muscle relaxation and/or which systemically reduce pulmonary capillary wedge pressure, and combinations of the foregoing.
- 36. A method according to Claim 21 wherein the disease is selected from the group consisting of fibromyalgia, gastric disorders and other systemic disorders, psychosis, other psychiatric disease, attention deficit disorder and systemic disorders, comprising vasospasm as a component.



- 37. A method according to Claim 21 wherein the disease is selected from the group consisting of systemic disorders comprising vasospasm as a component.
- 38. A titration system for diagnosing and treating a disease caused at least partially by insufficient cerebral perfusion, comprising in combination: operating a flow device to test for vasospasm, applying a dosage device which administers a vasospasm-reducing dosage of a medicine selected from the empirical group of medications which have the common characteristic of causing smooth muscle relaxation and/or which reduce pulmonary capillary wedge pressure, reoperating said flow device for testing over time and adjusting said dosage device to titrate said dosage to minimize occurrence and severity of said indications of vasospasm.

REMARKS

Any (small entity) charges required for the prosecution of this application should henceforth be charged to USPTO Deposit Account 20-0336 of Technology Licensing Co. LLC.

Please advise if anything further is required at this time.

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spectfully submitted

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